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Primary Fibril Nucleation of Aggregation Prone Tau Fragments PHF6 and PHF6*

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ABSTRACT: We performed replica exchange molecular dynamics and forward flux sampling simulations of hexapeptide VQIINK and VQIVYK systems, also known as, respectively, fragments PHF6* and PHF6 from the tau protein. Being a part of the microtubule binding region, these fragments are known to be aggregation prone, and at least one of them is a prerequisite for fibril formation of the tau protein. Using a coarse-grained force field, we establish the phase behavior of both fragments, and investigate the nucleation kinetics for the conversion into a β-sheet fibril. As the conversion is, in principle, a reversible process, we predict the rate constants for both the fibril formation and melting, and examine the corresponding mechanisms. Our simulations indicate that, while both fragments form disordered aggregates, only PHF6 is able to form β-sheet fibrils. This observation provides a possible explanation for the lack of available steric zipper crystal structures for PHF6*.

INTRODUCTION

Microtubule associated protein tau (MAPT), also known as tau, is a natively unfolded protein involved in the regulation of microtubule stability and therefore indirectly in the regulation of intracellular microtubule networks in the central nervous system. A repeat sequence, known as the microtubule binding region (MTBR), along with the neighboring proline-rich domain enables binding to microtubules. While tau is largely hydrophilic in nature and soluble, hydrophobic motifs in the MTBRs allow tau to form strong bonds with side chains of tubulin, resulting in an increased stability of the microtubule. Phosphorylation of tau lowers its affinity to bind to microtubules and therefore results in a loss of stability and rigidity, while vice versa microtubule stabilization occurs due to (partial) dephosphorylation of tau. While most tauopathies are due to mutations, hyperphosphorylation of tau is observed in many forms of dementia, causing a dramatic decrease in its binding affinity to microtubules. As a consequence, the resulting pool of dissociated tau assembles into β-like structures driven by hydrophobic interactions between the exposed MTBRs. First, dimers and small soluble oligomers form, which on their own display only little toxicity. As stacking continues, however, bundles of microtubules and therefore results in a loss of stability and rigidity, while vice versa microtubule stabilization occurs due to (partial) dephosphorylation of tau. While most tauopathies are due to mutations, hyperphosphorylation of tau is observed in many forms of dementia, causing a dramatic decrease in its binding affinity to microtubules. As a consequence, the resulting pool of dissociated tau assembles into β-like structures driven by hydrophobic interactions between the exposed MTBRs. First, dimers and small soluble oligomers form, which on their own display only little toxicity. As stacking continues, however, bundles of tau interact and settle into coiled-coil-like structures, so-called paired helical filaments (PHFs), with widths varying between 10 and 20 nm. Eventually, PHFs stick together, forming neurofibrillary tangles (NFT), one of the distinctive hallmarks of Alzheimer’s disease. However, while the extent of neurodegeneration correlates with the presence of NFTs, it is still not clear what specific process results in its neuronal toxicity and whether NFTs play a leading role in this mechanism. Gaining an understanding of tau–tau interplay and apprehending the properties resulting in tau assembly might lead to insights which can be used for the prevention of tau-induced toxicity in Alzheimer’s and other tauopathies.

Previous research highlights two particular fragments, VQIINK and VQIVYK in the MTBR repeats, respectively, called PHF6* and PHF6, as key sequences in tau fibrilization and assembly (see Figure 1). It is suggested that these motifs are notably prone to aggregate and that at least one of these hexapeptide sequences is a prerequisite for tau oligomer formation. Remarkably, only PHF6 seems to have an available steric zipper crystal structure. In 2001, Bergen et al. proposed two sequences containing these fragments as a subject for the study of tau assembly, R2/wt and R3/wt, which can be used to determine the effect that the PHF fragments have as part of a larger whole.

Molecular simulation can complement experimental work by giving detailed mechanistic insight into the aggregation process. However, simulations are hampered by several obstacles. First, system sizes are large, making the use of explicit force fields expensive. Second, the force fields themselves are not well suited for intrinsically disordered proteins. Third, the process of aggregation and fibrilization is a rare event that takes place on the minute to hour time scale in vitro and on the year time scale in vivo. Clearly, this puts stringent limitations on a straightforward all-atom approach.
We analyze the mechanisms, rates, and free energy landscapes, from coexistence, where the barrier to nucleation is substantial. and mechanism of sheet formation for temperatures not too far from the studied temperatures. In the Discussion section, we end with concluding remarks. We used the generic midresolution coarse-grained (CG) peptide model developed by Bereau and Deserno,29 which offers full sequence specificity. As this force field includes backbone hydrogen bonds, it has proven to be particularly accurate compared to other coarse-grained force fields when it comes to the prediction of the secondary structure of proteins. The model is parametrized such that both α-helical and β-sheet structures are equally accessible, which makes it more generic and realistic. This is partly achieved by including dipole interactions between carbonyl and amide groups. The solvent is treated implicitly in this model. For a detailed description of the model, we refer to ref 29. The integration was carried out by the ESPResSo31,32 MD package using Langevin dynamics with a 2 fs time step and a friction coefficient of 1.0 ps⁻¹. All MD snapshots were made with VMD.33

**METHODS**

**Force Field and Simulation Engine.** We used the generic midresolution coarse-grained (CG) peptide model developed by Bereau and Deserno,29 which offers full sequence specificity. As this force field includes backbone hydrogen bonds, it has proven to be particularly accurate compared to other coarse-grained force fields when it comes to the prediction of the secondary structure of proteins. The model is parametrized such that both α-helical and β-sheet structures are equally accessible, which makes it more generic and realistic. This is partly achieved by including dipole interactions between carbonyl and amide groups. The solvent is treated implicitly in this model. For a detailed description of the model, we refer to ref 29. The integration was carried out by the ESPResSo MD package using Langevin dynamics with a 2 fs time step and a friction coefficient of 1.0 ps⁻¹. All MD snapshots were made with VMD.33

**REMD Simulation.** Replica exchange molecular dynamics (REMD) simulates multiple system copies in parallel at different temperatures. Exchanging temperatures between replicas using a Monte Carlo algorithm allows these replicas to diffuse in temperature space, which can dramatically enhance sampling for complex systems with large free energy barriers.34 To determine the temperature range used for the REMD simulations, a trial simulation was initiated with a geometrical temperature distribution, after which we determined a theoretical exchange probability and calculated an optimized temperature distribution.35 This step is especially important when a (pseudo) phase transition is present, as the energy fluctuations at the phase transition prevent proper mixing of the replicas.

For the PHF6 and PHF6 dimerization, two fragments were positioned in a cubic box with edges of 51 Å, resulting in an overall concentration of 25 mM. REMD was performed using a temperature range of 210−518 K. The systems were simulated for 900 ns preceded by 100 ns equilibration, swapping replicas every 10 ps with an average exchange probability of ~30%.

To study aggregation of more than two peptides, we initialized three systems with N ∈ {8, 12, 20} peptides in a box with edges of 81 Å for N = 8 and 102 Å for both N = 12 and N = 20, corresponding to concentrations between 20 and 30 mM. REMD was performed with a temperature range of 210 to ~470 K in all cases. N = 8 and N = 12 were simulated for 900 ns, with an exchange of replicas every 20 ps, while N = 20 was simulated for 400 ns and exchanged replicas every 10 ps. All simulations were preceded by 100 ns equilibration and presented an exchange probability between 20 and 30%. The replica flow, j(Tᵢ), defines the capability of a system to move through temperature space and is defined as the fraction of replicas that diffused from the lowest to the highest temperature versus the temperature index i.35

Cluster analysis of these coarse-grained REMD simulations was in part carried out using home-written scripts.36 A peptide...
is part of a (fibrillar) cluster when four to six of its $C_n$ atoms are within 6.5 Å of the matching atom of any another peptide. The average interstrand distance is defined as the mean of distances between the centers of mass of all peptides.

The order of alignment in the system, represented by the nematic order parameter, $P_d$, is calculated using Wordom\textsuperscript{37,38} and is defined as

$$P_d = \frac{1}{N} \sum_{i=1}^{N} \frac{3}{2} \left( \hat{z}_i \cdot \hat{d} \right)^2 - \frac{1}{2}$$  \hspace{1cm} (1)

Here, $N$ is the number of peptides, $\hat{z}_i$ is a vector connecting the second and second-to-last $\alpha$-carbons of peptide $i$, and $\hat{d}$ is the director. The nematic order parameter can be employed to determine the overall order of alignment, as well as for the local positioning if only the alignment with a peptide's closest neighbor is considered. Values for the nematic order parameter range from $-\frac{1}{2}$ in the case of complete disorder, to a value of 1 for perfectly aligned peptides. In the case of a random distribution, $P_d = \sqrt{81/40 \pi N}$ describes the expected $P_d$ value for a system with $N$ peptides. To obtain information about transitions within a system, the weighted histogram analysis method (WHAM) is used to determine the system's average potential energy $U(T)$ as a function of temperature.\textsuperscript{22,39} From this energy, the heat capacity at constant volume, $C_V = \frac{\partial U}{\partial T}$, is calculated.

The free energy as a function of order parameter was computed from $F = -k_B T \ln \mathcal{P}(\bar{\lambda})$, where $k_B$ is the Boltzmann constant, $T$ the temperature, and $\mathcal{P}(\bar{\lambda})$ the normalized probability of finding a system in a certain configuration with order parameter $\bar{\lambda}$.

**Forward Flux Sampling.** The long time scales on which amyloid fibril formation occurs are related to high free energy barriers, making fibril nucleation a rare event. Enhanced sampling methods can address such rare events but require knowledge of the correct reaction coordinate which is often not available. The use of transition path sampling (TPS) alleviates this problem.\textsuperscript{40-44} Within the path sampling framework, the transition interface sampling (TIS) and the forward flux sampling (FFS) methods can efficiently compute rate constants.\textsuperscript{5,46} In TIS and FFS, the initial state $A$ and final state $B$ are separated in terms of an order parameter $\lambda$ such that $\lambda < \lambda_A$ in region $A$ and $\lambda > \lambda_B$ in region $B$. The boundaries of $A$ and $B$ are therefore defined by $\lambda_A$ and $\lambda_B$ (or $\lambda_0$ and $\lambda_\infty$ respectively). Between $A$ and $B$, several interfaces $i = 1, ..., n - 1$ are defined such that $\lambda_{i+1} > \lambda_i$. The expression for the rate constant $k_{AB}^{\lambda_B^{\lambda_A}}$ is given by

$$k_{AB}^{\lambda_B^{\lambda_A}} = \frac{\bar{\Phi}_A}{\bar{h}_B} P(\lambda_B | \lambda_A)$$  \hspace{1cm} (2)

where $\bar{\Phi}_A$ is the time-average of the effective positive flux of trajectories from $A$ that cross $\lambda_A$ and $\bar{h}_B$ is the average probability to be in $A$. The crossing probability $P(\lambda_B | \lambda_A)$ is the conditional probability that a trajectory from $A$ reaches $B$ before it returns to $A$, which can be expressed as a product

$$P(\lambda_B | \lambda_A) = \prod_i P(\lambda_{i+1} | \lambda_i)$$  \hspace{1cm} (3)

where $P(\lambda_{i+1} | \lambda_i)$ denotes the conditional crossing probability that a trajectory from $A$ that reaches $i$ also reaches $i + 1$, before returning to $A$. Here we employ the direct forward flux sampling (FFS) algorithm\textsuperscript{37} to study the primary fibril nucleation of the hexapeptides. The direct FFS algorithm is initiated with a straightforward MD run in region $A$, during which the order parameter $\lambda$ is followed over time.\textsuperscript{47} Each time the trajectory crosses boundary $\lambda_A$ in the positive direction, the configuration is stored until a total of $N_t$ configurations are generated. This procedure forms a set of starting configurations on $\lambda_A$ for trial runs toward the next interface $\lambda_B$ and simultaneously provides an estimate of the flux $\Phi_B = N_t/\tau$, where $\tau$ is the total length of the trial run. Next, the algorithm initiates several trial runs from a randomly selected starting configuration on interface $i$. If the trajectory reaches interface $\lambda_{i+1}$, the end-point is stored and the run is terminated. The trial run is also terminated if the order parameter returns to state $A$. Thus, each trajectory is free to recross previous interfaces before crossing the next. This procedure is repeated until $C_{n+1}$ configurations are generated on interface $i + 1$, and the number of runs needed to achieve this threshold is stored as $M_i$.

The FRESH software package is used to carry out the forward flux sampling simulations, as it complies well with ESPResSo.\textsuperscript{48} PHF6* and PHF6 were modeled using the peptideB force field. For all FFS simulations, $N = 12$ peptides were solvated in a box with edges of 102 Å, consistent with a concentration of ~20 mM. Water was implicitly modeled using Langevin dynamics with a main time step of 1 fs and a friction coefficient of 1.0 ps\textsuperscript{-1}.

Results from coarse-grained REMD simulations show that PHF6 self-assembles into two parallel $\beta$-sheets of six monomers; therefore, the order parameter is defined as the number of in-register $C_n$ contacts in the two largest clusters. If no clusters are present, the maximum number of in-register $C_n$ contacts between two single peptides is taken. Defining a proper distribution of order parameters is done on the basis of previous works on a similar subject,\textsuperscript{22} and is fine-tuned by a set of trial simulations. After calibration, five separate forward ($A \rightarrow B$) and five separate reversed simulations ($B \rightarrow A$) were performed using the settings shown in Table 1, and paused every 1 ps to calculate the order parameter. Here $i$ is the interface number, $C_i$ is the amount of paths collected up to that point, and $\lambda_i$ is the corresponding order parameter. To carry out the reversed FFS simulation, we define the auxiliary order parameter $\lambda'_i$ as $\lambda'_i = \lambda_i - \lambda$. The step-size $h$ is defined as the mean of distances $\lambda_i$.

### Table 1. Definition of Interfaces for FFS Simulations of the PHF6 and PHF6* Systems\textsuperscript{46}

<table>
<thead>
<tr>
<th>$i$</th>
<th>$C_i$</th>
<th>$\lambda_i$</th>
<th>$\lambda'_i$</th>
<th>$C_i$</th>
<th>$\lambda_i$</th>
<th>$\lambda'_i$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>200</td>
<td>8 (A)</td>
<td>8 (B)</td>
<td>30</td>
<td>4 (A)</td>
<td>4 (B)</td>
</tr>
<tr>
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<td>175</td>
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<td>12</td>
<td>29</td>
<td>8</td>
<td>8</td>
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<td>16</td>
<td>16</td>
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</tr>
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<td>27</td>
<td>16</td>
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<td>26</td>
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<td>20</td>
</tr>
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<tr>
<td>11</td>
<td>100</td>
<td>52 (B)</td>
<td>52 (A)</td>
<td>19</td>
<td>48 (B)</td>
<td>48 (A)</td>
</tr>
</tbody>
</table>

$^a$ $C_i$ is the number of target pathways. $\lambda_i$ denotes the $i$th interface value for the forward FFS run, $\lambda'_i$ for the reverse.
The simulations were carried out at a temperature of $T/(300 \text{ K}) = 1.138$ ($\sim 340 \text{ K}$), which is slightly below the ordering transition temperature, $T_0$. The value of $T_0$ could not be obtained for PHF6*, however, and a similar $T_0$ to that of PHF6 was used, namely, $T/(300 \text{ K}) = 1.12$ ($\sim 336 \text{ K}$).

The analysis of all FFS simulations was carried out using home-written scripts. If an FFS simulation is realized in both directions, i.e., $A \rightarrow B$ and $B \rightarrow A$, the free energy, $F(N_{C,i})$, for a certain order parameter $N_{C,i}$ can be extracted from the averaged simulation data as

$$F(N_{C,i}) = -k_B T \ln \rho(N_{C,i})$$

where $\rho(N_{C,i})$ is the probability to observe the number of order parameter $N_{C,i}$ atoms, and is given by

$$\rho(N_{C,i}) = \rho_A \Phi_{A,i} \tau_{A}(N_{C,i}; \lambda_A) + \rho_B \Phi_{B,i} \tau_{B}(N_{C,i}; \lambda_B)$$

where $\rho_A$ and $\rho_B$ are the steady state probabilities of being in $A$ and $B$, respectively, and are related via the detailed balance relation $\rho_A \Phi_{A,i} = \rho_B \Phi_{B,i}$. The flux out of either state $\Phi_{A,i}$ and $\Phi_{B,i}$ to the first interface is calculated during the initial straightforward MD simulation in the stable states. The function $\tau_{A}(N_{C,i}; \lambda_A)$ denotes the average time a system coming from either $A$ or $B$ has spent at order parameter $N_{C,i}$ and is, for the forward simulation, calculated as

$$\tau_{A}(N_{C,i}; \lambda_A) = \pi_A(N_{C,i}; \lambda_A) + \sum_{i=1}^{n-1} \pi_A(N_{C,i}; \lambda_A) \prod_{j=0}^{i-1} P(\lambda_j \rightarrow \lambda_{j+1})$$

where $\pi_A(N_{C,i}; \lambda_A)$ is the average time a run originating from interface $\lambda_A$ has spent at order parameter $N_{C,i}$. The function $\tau_{B}(N_{C,i}; \lambda_B)$ is calculated in a similar fashion with data from the reverse simulations.

To study the nucleation pathway, a cluster analysis is performed by examining the reactive pathways, i.e., all paths that reach interface $B$. For every snapshot, all clusters are classified, after which they are compared with the previous snapshot. The number of grown, shrunk, and unchanged clusters of size $n$ are registered in histograms $h_{\text{grow}}(n)$, $h_{\text{shrink}}(n)$, and $h_{\text{same}}(n)$, respectively. This process is repeated until every frame is compared with the one preceding it. The probability $P_x(n)$ of a cluster of size $n$ to belong to a certain category $x \in \{\text{grow, shrink, (remain the same)}\}$, $P_x(n)$, is

$$P_x(n) = \frac{h_x(n)}{h_{\text{grow}}(n) + h_{\text{shrink}}(n) + h_{\text{same}}(n)}$$

Sometimes it can be useful to compare only two of the three probabilities against each other. In that case, the value of the third histogram can be left out of the denominator.

**RESULTS**

**PHF6* and PHF6 Dimerization.** We performed REMD simulations of dimer systems of PHF6* and PHF6. The replica flow of both hexapeptides (see Figure 2) exhibits a good diffusivity of replicas through temperature space. The free energy landscapes as a function of the average interstrand distance for the two hexapeptides in Figure 3A and D show a nearly identical landscape for both PHF6* and PHF6. The free energy minimum at around 4 Å implies that the monomers are more likely to be in close proximity and form a dimer.

The free energy landscape of the nematic order parameter $P_2$, indicating the degree of alignment of the monomers in Figure 2A and E, is similar for both PHF6* and PHF6. The clear minimum at a relatively high nematic order parameter $P_2$ implies a proper alignment. Furthermore, Figure 3C and F shows that all six of the $C_6$ atoms are in-register with their counterpart, which coincides with an alignment in a parallel fashion.

**PHF6* and PHF6 Aggregation.** Next, we performed REMD simulations of the multiple peptide systems $N = 8$, $N = 12$, and $N = 20$. The replica flow for every system is shown in Figure 4. The PHF6* system shows an excellent replica flow for $N = 8$ and $N = 12$. Even though $N = 20$ deviates slightly from linear behavior, it shows a reasonably good replica flow, indicating sufficient diffusivity of replicas through temperature space.

A different situation occurs for PHF6, where $N = 12$ and $N = 20$ show a sigmoid-like curve. This indicates a much smaller although nonzero replica exchange ratio caused by a large difference in potential energy between systems on either side of the phase boundary. The $N = 8$ system, being a smaller system, still exhibits a reasonable flow.

The aggregation transition can be viewed as a (finite size) phase transition, and hence should have thermodynamic signatures such as a peak in the heat capacity. Figure 5 shows the calculated heat capacity for PHF6* and PHF6 with $N \in \{8, 12, 20\}$ peptides. PHF6* shows a peak at temperatures around 450 K, which corresponds to the peptide condensation peak, describing the transition from the gas (dilute phase) to the liquid (disordered oligomer phase). Other than that, no significant peaks are found for $N = 8$ and $N = 12$ peptides and only a small rise in $C_V$ can be observed around 300 K for $N = 20$, which might be an indication of a transition. Nevertheless, visual inspection of the configuration shows no strong sign of order. In contrast, for the PHF6 system, very distinct peaks occur for $N = 8, 12, 20$ peptides, akin to a phase transition. This peak occurs in a temperature range of 330 ± 20 K and corresponds to the transition between a disordered liquid at high temperatures and a systematic fibril-like oligomer arrangement at lower temperatures. For $N = 12$, the corresponding transition temperature $T_0$ is $T/(300 \text{ K}) = 1.143$ ($\sim 343 \text{ K}$). The change in transition temperature for different numbers of peptides is probably caused by the system size and/or poor replica flow.

Focusing on the $N = 12$ peptides, using a cluster and order parameter analysis, we compute the fraction of peptides that belong to an aligned ($\beta$-sheet) cluster. Upon an ordering...
transition, the aggregation fraction as a function of temperature exhibits an inflection point. In Figure 6, the PHF6* system shows an increasing aggregation fraction at lower temperatures, up to a value around 0.5 but not higher, as it fails to form an ordered fibril.22 This is in agreement with Figure 5 (left), in which also no transition is observed. In contrast, for the PHF6 system, the aggregation fraction as a function of temperature increases from zero for a disordered system at high temperatures to unity, indicating a fully ordered system at low temperature. The inflection point in the sigmoidal curve represents the transition temperature $T_0$ and is found at the same value that is suggested in Figure 5.

The free energy landscapes as a function of the total number of in-register $C_{\alpha}$ contacts for PHF6* and PHF6 are shown in Figure 7. In the case of PHF6*, a decrease in temperature goes hand in hand with a shift of the free energy minimum toward higher values of in-register contacts, which indicates a higher degree of organization at lower temperatures. The minimum observed at 210 K lies close to $N_{C_{\alpha}} = 35$, which is approximately half the theoretical maximum of ($11 \times 6 = 66$). This behavior is known for substances which polymerize at lower temperature and form so-called cluster liquids.22 In such liquids, small clusters are observed but no fibrils are formed.
Even though PHF6 does not reach the theoretical maximum of 66 in-register contacts either, it shows a significantly different behavior than PHF6*. A free energy minimum at values as high as 60 in-register contacts at a temperature $T/(300 \text{ K}) = 0.7$ ($\sim 210 \text{ K}$) suggests a degree of order consistent with full fibrilization.

Figure 8 shows the normalized cluster size distributions $\rho(n)$. In the case of PHF6*, the exponential decrease in probability for increasing cluster sizes indicates that no fibrilization takes place. At low temperature (purple line), dimers, trimers, and tetramers are present, indicating the existence of a polymerized or cluster liquid.22

In the case of PHF6, the clear peak at $n = 6$ monomers implies that this peptide forms stable aggregates and suggests that fibrils consist of two sheets of six peptides. We found no evidence for the formation of different combinations of sheets, such as a single $\beta$-sheet consisting of 12 monomers, or a pentamer/heptamer sheet combination, an octamer/tetramer...
sheet combination, or even the combination of three tetramer sheets.

Figure 9 shows the free energy landscape as a function of the overall degree of alignment of the system using the nematic order parameter $P_2$. The free energy minima of PHF6* remain the same for all temperatures, which coincides with the expected value for a randomly ordered system where $N = 12$, as $A_{\parallel} = \sqrt{\frac{81}{40\pi12}} \approx 0.23$, confirming again PHF6*'s incapacity to form fibrils. Figure 9 (right) shows a similar free energy curve at the same location for PHF6 at high temperatures where the oligomer is disordered. At lower temperatures, two free energy minima are observed. The minimum at $P_2 \approx 0.5$ corresponds to a state in which the monomers of the first sheet are not perfectly aligned with the monomers of the second sheet, whereas the second minimum, at 0.95, belongs to a configuration where all monomers are well-aligned. Visual inspection of the simulations shows a parallel alignment of the monomers in each sheet, whereas the sheets are oriented in opposing directions.

**Primary Nucleation Kinetics of PHF6.** We performed FFS simulations at a temperature of $T/(300 K) = 1.138 (~340)$ K for the PHF6 systems, starting at an initial disordered liquid state A and ending at the final fibril configuration B. Table 2 shows an overview of the resulting crossing probabilities between each set of neighboring interfaces along with the average path length, $\bar{l}$, for the FFS simulation of PHF6. The crossing probability is relatively constant (around 0.2) up to $\lambda_{\parallel}$ corresponding to 32 in-register contacts, indicating a steep barrier. At higher values, the crossing probability steadily increases as a function of $\lambda$, indicating that the barrier becomes less steep.

The reversed simulation ($B \rightarrow A$, dissolution of the fibril) shows a relatively constant crossing probability of $30 \pm 10\%$. The qualitative difference in crossing probability behavior between forward and reverse FFS simulations suggests some degree of hysteresis; i.e., the association and dissociation processes follow different pathways.

The flux, $\Phi$, and rate constants, $k$, are also reported in Table 2. Clearly, self-assembly of PHF6 fibrils proceeds faster than its dissociation, with the forward reaction rate $k_B = 6.7 \times 10^9$ s$^{-1}$ more than 10 times higher than the reverse reaction rate $k_A = 5.9 \times 10^8$ s$^{-1}$. The equilibrium constant is $K = k_B/k_A = 10.29$ corresponding to a difference in free energy between the two states, $\Delta F = -k_B T \ln K = -2.43 k_B T$. This negative value of $\Delta F$ confirms the system’s tendency to form fibrils at this temperature. The corresponding equilibrium (steady state) population, $\rho_B = \frac{k_B}{k_B + k_A} = 0.919$, indicates the probability of finding the system in state B, which is much larger than the probability of finding the system in state A, namely, $0.081$ as $\rho_B + \rho_A \approx 1$ at this temperature.

Figure 10 shows the (partial) free energy landscape for the forward $A \rightarrow B$ as well as the reversed $B \rightarrow A$ simulation. The free energy curves fluctuate near the borders due to poor sampling of the fully dissociated and associated state during the forward and backward simulations, respectively.

Combining the forward and backward free energies and weighting them properly by applying eq 4 results in the overall free energy landscape as a function of the number of in-register contacts, which is also depicted in Figure 10. Apart from several small metastable minima, one distinct free energy barrier is observed near 30 in-register contacts. This value is roughly in agreement with the point where the forward crossing probabilities start to increase. The middle inset of Figure 10 (right) shows a transition structure from the forward paths, consisting of several half formed $\beta$-sheets. This structure is just one of the possible transition states that can be seen as critical nuclei in the primary nucleation process of fibril formation. Such structures have relatively high free energy as the peptides have lost configurational entropy but have not yet gained sufficient stabilization from favorable interactions, such as the

![Figure 9](image)

Figure 9. Free energy vs the nematic order parameter $P_2$ for $N = 12$ peptides of PHF6* (left) and PHF6 (right).

<table>
<thead>
<tr>
<th>$i$</th>
<th>$P(x_i, i)$</th>
<th>$l_{\text{forward}}$ (ps)</th>
<th>$P(x_i, i')$</th>
<th>$l_{\text{backward}}$ (ps)</th>
</tr>
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<tr>
<td>1</td>
<td>0.1871</td>
<td>5</td>
<td>0.2406</td>
<td>4</td>
</tr>
<tr>
<td>2</td>
<td>0.1852</td>
<td>6</td>
<td>0.2045</td>
<td>3</td>
</tr>
<tr>
<td>3</td>
<td>0.1846</td>
<td>15</td>
<td>0.2996</td>
<td>8</td>
</tr>
<tr>
<td>4</td>
<td>0.1999</td>
<td>52</td>
<td>0.2957</td>
<td>19</td>
</tr>
<tr>
<td>5</td>
<td>0.2017</td>
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<td>0.2604</td>
<td>49</td>
</tr>
<tr>
<td>6</td>
<td>0.2562</td>
<td>153</td>
<td>0.3105</td>
<td>81</td>
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<tr>
<td>7</td>
<td>0.4178</td>
<td>207</td>
<td>0.2700</td>
<td>204</td>
</tr>
<tr>
<td>8</td>
<td>0.6024</td>
<td>399</td>
<td>0.2467</td>
<td>446</td>
</tr>
<tr>
<td>9</td>
<td>0.6680</td>
<td>886</td>
<td>0.3724</td>
<td>476</td>
</tr>
<tr>
<td>10</td>
<td>0.8326</td>
<td>1222</td>
<td>0.3940</td>
<td>784</td>
</tr>
<tr>
<td>$B$</td>
<td>0.9188</td>
<td>1645</td>
<td>0.3803</td>
<td>1344</td>
</tr>
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</table>

$\Phi \approx 5.5 \times 10^7$ s$^{-1}$ $l_{\text{forward}} = 6.2 \times 10^9$ s$^{-1}$ $l_{\text{backward}} = 5.9 \times 10^8$ s$^{-1}$
formation of the second sheet. The free energy difference between state A and state B of $\sim 2 \ k_B T$ approximates the calculated free energy difference of 2.43 $k_B T$. This result is in agreement with the REMD predictions in Figure 7(right).

The transition path ensemble (TPE) of the fibril formation is available from the FFS simulation. We analyze these paths to obtain insight into the growth kinetics of fibril-like structures in the oligomer. The kinetic behavior of the peptide clusters in the oligomer along the TPE is shown in Figure 11, which compares the growth probability, $P_{\text{grow}}$, with the shrink probability, $P_{\text{shrink}}$, such that the sum adds up to unity. Clusters consisting of up to four monomers have the tendency to grow, whereas larger clusters are more likely to reduce in size. This lack of growth is most likely due to depletion of monomers in the droplet. In addition, it appears to be more favorable for a monomer to dimerize and start forming a second sheet than to anchor to an existing cluster of four. Figure 11 (right) includes the probability that a cluster remains of the same size, which de facto is a measure of an oligomer’s stability. The probability $P_{\text{same}}$ increases up until $n = 6$, after which the stability declines. This behavior agrees with the observation that two sheets of six peptides are formed in the coarse-grained REMD simulations. Clusters containing more than eight monomers seem to shrink at all times, implying instability.

We also performed an FFS simulation of PHF6*. The crossing probabilities are reported in Table 3 and are remarkably low, and become basically zero at $\lambda_{9y}$ corresponding to 40 in-register contacts. The relatively short transition pathways at higher $\lambda$’s ($\sim 10\%$ of the path length observed for PHF6) indicate that the encountered states are coincidental structures and do not correspond to a viable transition path. The computed rate for this process is lower than $10^{-10} \text{ s}^{-1}$, corresponding to once per every few centuries. These

Table 3. Crossing Probability for the FFS Simulation of PHF6* along with the Average Path Length ($\bar{l}$)

<table>
<thead>
<tr>
<th>$\lambda$</th>
<th>$P_{\text{grow}}$</th>
<th>$P_{\text{shrink}}$</th>
<th>$\bar{l}_{\text{forward}}$ (ps)</th>
<th>$\bar{l}_{\text{backward}}$ (ps)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.5912</td>
<td>0.4801</td>
<td>2</td>
<td>3</td>
</tr>
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<td>2</td>
<td>0.0111</td>
<td>0.3742</td>
<td>3</td>
<td>7</td>
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<td>3</td>
<td>0.0126</td>
<td>0.5513</td>
<td>7</td>
<td>13</td>
</tr>
<tr>
<td>4</td>
<td>0.0022</td>
<td>0.6042</td>
<td>13</td>
<td>35</td>
</tr>
<tr>
<td>5</td>
<td>0.0094</td>
<td>0.6653</td>
<td>35</td>
<td>69</td>
</tr>
<tr>
<td>6</td>
<td>0.0016</td>
<td>0.7871</td>
<td>69</td>
<td>137</td>
</tr>
<tr>
<td>7</td>
<td>0.0239</td>
<td>0.7675</td>
<td>137</td>
<td>223</td>
</tr>
<tr>
<td>8</td>
<td>0.0033</td>
<td>0.7341</td>
<td>223</td>
<td>456</td>
</tr>
<tr>
<td>9</td>
<td>5.2912 $\times$ 10^{-5}</td>
<td>0.9338</td>
<td>456</td>
<td>903</td>
</tr>
<tr>
<td>10</td>
<td>n/a</td>
<td>0.9348</td>
<td>903</td>
<td>1800</td>
</tr>
<tr>
<td>11</td>
<td>n/a</td>
<td>0.9524</td>
<td>1800</td>
<td>3600</td>
</tr>
</tbody>
</table>

$\Phi$ $3.6 \times 10^{10} \text{ s}^{-1}$ $7.0 \times 10^{10} \text{ s}^{-1}$

$k$ $1.55 \times 10^{10} \text{ s}^{-1}$ $8.4 \times 10^{8} \text{ s}^{-1}$
observations support the hypothesis that \( \text{PHF6}^* \) is unable to form stable \( \beta \)-sheet-based fibrillar aggregates. To test this hypothesis further, we initiated a reverse FFS simulation, starting from a fibril structure taken from the \( \text{PHF6} \) FFS simulations. The reverse FFS simulations showed that this structure was not stable, and fell apart very quickly. Indeed, the dissociation rate in Table 3 is 4 orders of magnitude higher than that of \( \text{PHF6} \), and at least 17 orders of magnitude faster than the nucleation rate of \( \text{PHF6}^* \).

\section*{Discussion}

While the dimer properties of \( \text{PHF6} \) and \( \text{PHF6}^* \) are almost identical, when looking at the tendency of \( \text{PHF6}^* \) and \( \text{PHF6} \) to self-assemble into fibril-like structures, a remarkable difference is encountered. The simulations show that \( \text{PHF6}^* \) is not able to form stable aggregates, which unfortunately cannot be straightforwardly compared to the literature, as no crystal structure has yet been reported for this peptide.\(^{50}\) In contrast, \( \text{PHF6} \) appears to be very well able to form fibrils. Our results can be compared to the study by Li et al.,\(^{25}\) who performed Monte Carlo simulations of the \( \text{PHF6} \) fragment employing an implicit solvent all-atom model. They found evidence for a nucleation process, similar to our results. However, as their simulation was done at lower temperature, the authors report a very fast aggregation and observed hardly any nucleation barrier. For a system of 12 peptides, they observed the formation of stable oligomers consisting of two sheets held together by hydrophobic interactions. In contrast to our findings, initially these sheets contain a mixture of parallel and anti parallel strands, which have to reorder in order to initiate further fibril growth.

The higher aggregation propensity of \( \text{PHF6} \) with respect to \( \text{PHF6}^* \) is in accordance with numerous studies.\(^{13,18,19,21,51,52}\) The difference can be accounted for by looking at the difference in amino acid sequence and in their corresponding interaction coefficients, Table 4. Here \( \epsilon'_i \) is an averaged parameter determining the strength of interaction between specific residues. The main difference between the two peptides lies in the hydrophobic triad, VIX, where X is asparagine for \( \text{PHF6}^* \) and tyrosine for \( \text{PHF6} \). Asparagine has a significantly lower interaction coefficient than tyrosine, resulting in an ~40% lower interaction with the other residues, which apparently can make the difference between aggregating or not.

Experiments show that eliminating \( \text{PHF6}^* \) from \( \text{R2/wt} \) does still lead to some tau aggregation, while removing \( \text{PHF6} \) prevents all aggregation.\(^{19}\) However, \( \text{PHF6}^* \), which is part of \( \text{R2/wt} \), does not fibrilize, whereas \( \text{PHF6} \) does form fibrils. This observation is rather counterintuitive and seems to contradict the observation that \( \text{R2/wt} \) is more aggregation prone than \( \text{R3/wt} \).

The disability of \( \text{PHF6}^* \)'s to form a stable fibril under the peptideB force field is confirmed by the results obtained from the FFS simulations. As mentioned before, we are unable to compare this to experiments, as no crystal structure has been reported for this peptide.\(^{50}\) However, the fact that our simulations seem to predict that this hexapeptide is not able to form fibrils might explain this lack of crystal structure.

The \( \text{PHF6 N} = 12 \) system forms fibril nuclei consisting of two \( \beta \)-sheets of six peptides in case. In these sheets, monomers are stacked in a parallel fashion, whereas the two sheets lie in an antiparallel fashion. This structure has also been experimentally observed for this hexapeptide and is described as class 1 in the eight classes of steric zippers, which describes the fibril organization of aggregation prone peptides.\(^{50,53,54}\)

On the basis of visual inspection and cluster analysis, we can propose a nucleation mechanism, as visualized in Figure 12. Starting from a liquid of unorganized monomers (i), a single \( \beta \)-sheet is formed consisting on average of four monomers (ii). Even though occasionally more peptides lock onto the tetramer and enlarge the existing sheet (iii), the most likely next step in the process involves the formation of a second \( \beta \)-sheet (iv). Note that in most cases the two sheets form almost simultaneously or very rapidly succeeding each other. The next step is the alignment of the two existing sheets (v). The overall barrier seems to have been reached in the last step, where the remaining peptides anchor to the existing oligomer and form a fibril-like structure (vi).

While the reverse process in principle should follow the same mechanism due to the detailed balance, a certain amount of

\[
\begin{array}{cccccc}
\epsilon'_i (\text{PHF6}^*) & 5.43 & 1.74 & 6.79 & 6.79 & 1.54 & 0.99 \\
\text{residue} & V & Q & I & \bar{V} & N & K \\
\epsilon'_i (\text{PHF6}) & 5.43 & 1.74 & 6.79 & 5.43 & 4.29 & 0.99 \\
\end{array}
\]

**Figure 12.** Schematic representation of the \( \text{PHF6} \) nucleation mechanism inferred from the forward FFS paths.
hysteresis can be expected. We therefore also inspected the dissociation process in the reverse FFS path ensemble (for a visual representation, see Figure 13). In the dissociation process, the peptides of the formed fibril (i) first melt at the ends of the fibril (ii) and rearrange (iii). Then, one of the β-sheets melts first (iv), after which the second one melts (v). There are many similarities with the nucleation pathways, such as the presence of the 1-β-sheet intermediate state, but there are also striking differences. For instance, the locking transition of the two β-sheets observed in the nucleation process does not occur during melting. Notwithstanding these differences, the reverse mechanism is in agreement with the nucleation pathways, showing that the FFS sampled more or less the correct path ensemble.

These simulations confirm that this peptide nucleates via a two-step nucleation (2SN), where the first step is the appearance of nonfibrillar disordered oligomers in the solution after which they convert into fibrils, in agreement with Ostwald’s step rule. Indeed, 2SN has been known to occur for peptides with a strong hydrophobic nature.\textsuperscript{22,36,55}

The barrier of 8 \( k_B T \) predicted by the FFS simulations in Figure 10 seems in reasonable agreement with the fibrilization rate of 6.70 \( \times 10^3 \) s\(^{-1}\), assuming a transition state theory prefactor on the order of 10\(^9\) s\(^{-1}\). Any underestimation of the barrier is probably caused by plotting the free energy as a function of an approximate order parameter, and the presence of multiple pathways. In fact, the predicted rate of 6.70 \( \times 10^3 \) s\(^{-1}\) on itself is a very high rate for a process that in the central nervous system occurs on a time scale of months to years. This overestimation of rate can be attributed to several reasons. First, molecular dynamics in the CG force field is faster compared to all-atom force fields with an explicit solvent, due to their softer effective potentials and the use of an implicit solvent. Second, the simulations are done at a relatively high concentration. Indeed, at such high concentrations, even straightforward MD already shows some spontaneous aggregation within a few hundred nanoseconds.\textsuperscript{54} The concentration of intracellular \( \tau \)-actin lies around 3 \muM, the majority of which (>99%) is tightly bound to the microtubule.\textsuperscript{6,57} Even though the free \( \tau \)-actin concentration is elevated in patients with Alzheimer’s disease or other forms of dementia, the concentrations used for this study are much higher than those in the human brain.\textsuperscript{58–60} Moreover, no other molecules were included that might interfere or slow down the assembly of PHF6 in the brain.

\section*{CONCLUSIONS}

Experimental work has shown that PHF6* and PHF6 are important sequences for \( \tau \)-actin to aggregate.\textsuperscript{19} In our simulations, we observe that PHF6* and PHF6 dimerize, as do the R2/wt and R3/wt sequences. On the other hand, while both hexapeptides aggregate into a disordered oligomer, only the PHF6 undergoes a two-step nucleation and converts into a fibrillar structure consisting of two parallel β-sheets. The PHF6* peptide systems do not show any sign of fibril formation in our REMD simulations but remain in a disordered polymerized liquid state. It follows that, while PHF6* could assist in the aggregation of \( \tau \)-actin, the hexapeptide itself is not interacting strongly enough to induce fibrilization.

The kinetics of PHF6* and PHF6 conversion from a disordered liquid oligomer into a fibril-like structure was investigated using forward flux sampling simulations. The PHF6 peptide forms fibrils with a rate constant of \( k_{AB} = 6.70 \times 10^3 \) s\(^{-1}\) at a temperature of 340 K, whereas the PHF6* did not fibrillize at all. Reverse FFS simulations confirmed that a PHF6* fibril is unstable and quickly falls apart.

The results in this work give qualitative insight into the primary nucleation kinetics of short fragments of the tau protein. The stability and kinetic predictions should be taken qualitatively, as the force fields are approximate. Moreover, the use of a CG force field will render the predicted kinetics too fast, due to smoothing of the potentials. Nevertheless, our results seem to indicate a physical reason for the fact that a PHF6* steric zipper crystal structure is not available. These results are also in agreement with the experimental observation that, while PHF6 is a prerequisite sequence for \( \tau \)-actin aggregation, PHF6* merely enhances aggregation.\textsuperscript{19}

In future work, we will investigate the fibrilization of R2/wt and R3/wt with the CG models. In addition, we will improve the predictions of this work by using full atomistic models to access the kinetics of the hexapeptides.

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The authors declare no competing financial interest.

## REFERENCES


